PATENT SPECIFICATION

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COMPLETE SPECIFICATION

NO DRAWINGS

Process for the Preparation of Emulsifiers for Ointment Base Compositions

We, DEHYDAG DEUTSCHE HYDRIERWERK GmbH., a German Company, of 67, Henkelstrasse, Dusseldorf, Germany, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:—

This invention relates to a process for the 10 production of emulsifiers for ointment bases.

The present invention provides a composition for an ointment base which contains a mixed ester from a pentaerythritol-di-fatty acid ester and a citric acid-di-fatty alcohol seter in the molar ratio 1:1. These compositions are distinguished by being odourless and having a high, steady water-binding capacity.

It has been found that specially valuable 20 products with the above-described valuable properties are obtained when mixed ester products which possess lipophilic residues simultaneously with both the polybasic citric acid residue and the polyhydric alcohol 25 residue are prepared by an esterification process under an inert gas. Those products are particularly advantageous which are derived from pentaerythritol as the polyhydric alcohol.

The particular technical value of the compositions prepared according to the process of the invention rests not only on their structure and the advantages resulting therefrom, but also on the ease of production. It is known that the preparation of citric acid esters by the usual esterification methods causes difficulties, since the citric acid on relatively long heating is converted partly into unsaturated aconitic acid with splitting 40 off of water which leads to troublesome resinous compounds in the further course of the reaction.

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made that these difficulties may be avoided if aqueous citric acid is used for the 45 esterification. The use of an aqueous component in the esterification proless is entirely contrary to the usual customs, since it normally causes a delay in the course of the reaction and therefore favours side reactions. The 50 esterification is carried out by adding to the given fatty alcohol, heated at 140-170°C., for the most part at 160°C., and under a pressure of about 25 mm Hg, only as much aqueous citric acid as reacts simultaneously 55 with the alcohol. Every side reaction and consequently resinification is prevented by the rapid esterification and a satisfactory esterification product light-coloured obtained.

The preparation of the second esterification product, the pentaerythritol di-fatty acid ester, takes place by the usual process, in which case preference is given to the production by inter-esterification of a fatty 65 acid methyl ester, since in this way especially light-coloured products with an acid value below 1 are obtained.

Examples of the preparation of the pentaerythritol di-fatty acid ester include:—

(i) 136 kg pentaerythritol with 470 kg. coconut fatty acid methyl ester and 1.8 kg. sodium methylate as esterification catalyst are placed in an agitator.

The contents of the apparatus are highly 75 heated with stirring. At approximately 70°C. methyl alcohol is distilled from the reaction mixture. When no further methyl alcohol is distilled over the final residues of alcohol is removed in vacuo. After 2 hours the acid 80 number has dropped below 2 and the esterification is concluded.

(ii) 100 kg coconut fatty acid are employed. With stirring there are added; 31 kg pentaerythritol, 200 kg 50% soda lye as 85 esterification catalyst. Esterification is

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effected for one hour at 180°C, and for a further 5 hours at 180°-190°C, at a pressure of the esterification can be followed by the of approximately 50 mms. During this time water of reaction which passes over. The a weak flow of nitrogen is passed through ester is light in colour, has an esterification 5 the liquid by means of a dip tube serving value of 241 and an acid value of 80 to 85, 70 as a boiling capillary. The esterification is as is necessary for the further preparation concluded when the acid number of the reof the mixed condensate. action mixture has dropped below 2. Preparation of a pentaerythritol di-coconut oil fatty acid ester. The interesterification of the above two 10 esterification products is carried out in an 76 parts by weight of pentaerythritol are 75 inert atmosphere and is concluded when the reacted with 260 parts by weight of coconut acid value of the reaction mixture falls below one. This interesterification process oil fatty acid methyl ester with a saponification value of 240 and 1 part by weight sodium methylate as esterification may be carried out at reduced pressure and 15 elevated temperature and preferred values catalyst until the acid value has fallen below 80 for the pressure and temperature are 25 mms. 2. The pentaerythritol di-coconut oil fatty at Hg and 180° to 190°C., respectively. acid ester formed is freed from the accom-Catalysts are not generally required.

Accordingly the present invention propanying soap with 1% of fuller's earth.
(c) Preparation of the citric acid mixed 20 vides a process for the preparation of emulsicondensate. fiers for ointment bases in which a mixed 140 parts by weight of citric acid dioctaester is prepared by an interesterification decyl ester from (a) and 115 parts by weight of pentaerythritol di-coconut oil fatty acid process under an inert gas from a citric aciddi-fatty alcohol ester prepared by esterifiester from (b) are esterified in an atmosphere of inert gas at a pressure of 25 mms of Hg 90 25 cation of a fatty alcohol with aqueous citric acid and a pentaerythritol-di-fatty acid ester and a temperature in the range 180°-190°C. in the molar ratio of 1:1. until the acid value of the reaction product The citric acid mixed esters obtainable lies below 1. After the esterification is finished, the mixed ester is bleached with using the process of the invention from a 0.1% (referred to the total weight) of 40% 95 30 pentaerythritol-di-fatty acid ester and a citric hydrogen peroxide. A mixed ester of waxacid-di-fatty alcohol ester display very valuable properties. The stability to temperature yellow colour with an acid value below 1, of the emulsions produced therewith is outa saponification value of 224 and a hydroxyl standing, the limit of temperature being 35 especially high, at about 50°C. The watervalue of 75 is obtained. 100 (d) Preparation of an ointment base. binding capacity is likewise considerably 35 parts by weight of vaseline better and not only are relatively large 30 " decyloleate 25 37 ... amounts of water absorbed substantially " cetyl alcohol " ozocerite, white, 70/72° more rapidly, but the larger total amount of ** ** " hard paraffin, 50/52° 105 40 water may also be incorporated in larger portions. In addition, the ointments obtained using the usual qualities of Vaseline " paraffin, viscous using the usual qualities of Vaseline (Registered Trade Mark) are pure white and , aluminium stearate with addition of 12 parts by weight of the no longer yellow as hitherto, and are citric acid mixed ester according to (c) above 45 furthermore practically odourless and easily and durably perfumed. A further important are melted together on the water bath, 110 stirred until homogeneous and allowed to advantage is that they are easier to preserve cool. An ointment base is obtained which and have far better keeping quality on posesses a high water absorption capacity. fairly long storage. The ointments obtained After suitably perfuming or after addition 50 also display a smoother softer structure, a of pharmaceutically active substances, the 115 more pliable consistency, i.e. they do not ointment base may be used as such. But stick, provide a subjectively more pleasant there may also be incorporated in the ointimpression on the skin and are absorbed ment base up to three times its amount of more easily by the skin. The ointments prewater, when salves (water in oil emulsions) of various consistency are obtained. The 120 amount of water used depends upon the 55 pared with the products according to the invention are dermatologically completely special purpose for which the salve is to satisfactory. (a) Preparation of a citric be used and may be adjusted to any desired Example 1. acid dioctadecyl ester. 115 parts by weight of octadecyl alcohol Example 2. of hydroxyl number 206 and 54 parts by Mixed esters from the following pairs of esters may be prepared similarly: citric acid dilauryl ester and pentaerythritol-distearic weight of aqueous citric acid (consisting of 44 kg of citric acid dissolved in 10 kg of acid ester, citric acid-dilauryl ester and water) are reacted so that only as much pentaerythritol-dicoconut oil fatty acid ester, 130 65 aqueous citric acid is added as can react with

the given alcohol at one time. The course

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citric acid-di-Guerbet alcohol (C₂₂) ester and pentaerythritol-distearic acid ester, citric acid-dioctadecanediol ester and acid-dioctadecanediol ester and penta-erythritol-di-coconut oil fatty acid ester, 5 citric acid-dioctadecanediol ester and pentaerythritol-distearic acid ester, citric aciddidodecyl ester and pentaerythritol-dicoco-

nut oil fatty acid ester.

Among all the citric acid mixed esters 10 specified, the mixed ester from citric aciddioctadecyl (stearyl) ester and penta-erythritol-di-coconut oil fatty acid ester is most suitable for use as an emulsifier in ointment bases.

WHAT WE CLAIM IS:-

1. A process for the preparation of emulsifiers for ointment bases in which a mixed ester is prepared by an esterification process under an inert gas from a citric acid-20 di-fatty alcohol ester, prepared by esterifi-cation of a fatty alcohol with aqueous citric

acid, and a pentaerythritol-di-fatty acid ester in the molar ratio of 1:1.

2. A process as claimed in claim 1 in which the citric acid-di-fatty alcohol ester 25 is citric acid-dioctadecyl (stearyl) ester.

3. A process as claimed in any preceding claim in which the pentaerythritol-di-fatty acid ester is pentaerythritol di-coconut oil fatty acid ester.

4. A process for the preparation of ointment bases substantially as hereinbefore described with reference to the Examples.

5. Emulsifiers for ointment bases when prepared by the process of any one of claims 35

6. Ointments containing the emulsifiers claimed in claim 5.

> W. P. THOMPSON & Co., 12, Church Street, Liverpool, 1. Chartered Patent Agents.

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